

Eur päisches Patentamt
Eur pean Patent Office
Offic européen d s brevets



11 Publication number:

0 647 450 A1

(12)

# **EUROPEAN PATENT APPLICATION**

21 Application number: 93114475.2

(1) Int. Cl. 6: A61K 47/48

Date of filing: 09.09.93

② Date of publication of application: 12.04.95 Bulletin 95/15

Designated Contracting States:
 AT BE CH DE DK ES FR GB GR IE IT LI LU NL
PT SE

Applicant: BEHRINGWERKE Aktlengesellschaft Postfach 1140 D-35001 Marburg (DE)

2 Inventor: Bosslet, Klaus, Dr.
An der Haustatt 64
D-35007 Marburg (DE)
Inventor: Czech, Jörg, Dr.
Kreutzackerweg 2a
D-35041 Marburg (DE)
Inventor: Hoffmann, Dieter, Dr.
Feuerdornweg 123
D-35041 Marburg (DE)

Inventor: Tillequin, François, Prof. 70, rue de l'Amiral Mouches

F-75013 Paris (FR)

Inventor: Florent, Jean-Claude, Dr.

23 rue des Causus F-91940 Les Ulis (FR)

Inventor: Azoulay, Michel, Dr.

16 rue du Regard F-75006 Paris (FR)

Inventor: Monneret, Claude, Dr.

9 avenue Lamoricière F-75012 Paris (FR)

Inventor: Jacque, Jean-Claude, Prof.

46 rue du Planty

F-86360 Buxerolles (FR)

Inventor: Gesson, Jean-Pierre, Prof.

"La Germonière"-Montassie

F-86360 Chansseneuilo du Poltou (FR)

Inventor: Koch, Michel, Prof.

116 Elysées 2

F-78170 La Celle Saint Cloud (FR)

Inventor: Vasella, Andrea A., Prof.

Langackerstrasse 7 CH-8057 Zürich (CH) Inventor: Hoos, Roland Affolternstrasse 107 CH-8050 Zürich (CH)

- Improved prodrugs for enzyme mediated activation.
- © Enzymatically cleavable prodrugs with reduced Michaelis-Menten constant (Km) are described.

This invention r fers to enzymatically cleavable prodrugs with reduced Michaelis-Ment n constant (Km). A prodrug may be defined as a chemical which is non-toxic and pharmacodynamically inert, but which can b transformed in vivo to a pharmacologically active drug.

The inv ntion refers to the field of drug-targeting, which deals with site-specific delivery of drugs in vivo. Site-specific delivery preferably increases the selectivity of drugs and reduces their undesirable side effects.

One potential approach to achieve a site-specific delivery consists in applying untoxic prodrugs which can be site-specifically activated to cytotoxic drugs using prelocalized prodrug cleaving catalysts like enzymes, muteins derived from enzymes, catalytic antibodies, antibody enzyme conjugates or fusion proteins.

This approach combines the advantage of drug delivery via prodrugs (i.e. increased stability, adjusted solubility, improved route of administration, more favourable distribution, improved pharmacokinetics, bypassing resistance; T.A. Connors, Xenobiotica 16, 975-988, 1986) with the preferential tumour specific activation mediated by a catalytic principle. The use of exogenous enzymes or polyclonal antibody enzyme conjugates for prodrug activation was pioneered by Graffi (Deutsche Offenlegungsschrift 22 12 014), and Philpott et al. (J. Immunol. 111, 921, 1973).

More recently the original teaching from Graffi and Philpott was exemplified and improved by the use of monoclonal antibody enzyme conjugates as prodrug activating catalysts (Bagshawe et al., Brit. J. Cancer, 58, 700, 1988; Senter et al., Bioconjugate Chem. 4, 3-9, 1993) or fusion proteins (Bosslet et al., Brit. J. Cancer, 65, 234-238, 1992; Goshorn et al., Cancer Res. 53, 2123-2127, 1993).

Despite these improvements, the systems described so far have some major disadvantages for clinical applications:

- a) monoclonal antibody enzyme conjugates produced by chemical coupling have as a major drawback a strong immunogenicity in man due to the xenogenic origin of the antibody moiety and the enzyme (Bagshawe et al., Disease Markers 9: 233-238, 1991). As a consequence of this high immunogenicity repetitive applications in man are possible only to a very limited extent;
- b) fusion proteins consisting of non-humanised binding moieties and xenogenic enzymes produced by recombinant DNA technology will be immunogenic in man as well with disadvantages comparable to monoclonal antibody enzyme conjugates, if repetitive applications are needed;
- c) fusion proteins consisting of humanized binding moieties and human enzymes will probably not be very immunogenic in man most probably allowing repetitive treatment cycles in man. Nevertheless, the two major disadvantages of human fusion proteins are the possibly lower turnover rate (Vmax) of the human enzyme moiety as well as the possibly higher prodrug (substrate) concentration needed to obtain significant catalysis in comparison to xenogenic enzymes having a high turnover rate and a low Michaelis-Menten constant (Km).

This limitation of human fusion proteins (low Vmax and high Km) given by the intrinsic nature of the human enzyme moiety can be overcome by state of the art methodology only to a very limited extent (factor 4) by random mutagenesis in the active site of the enzyme (Munir et al., PNAS USA 90:4012-4016, 1993).

Surprisingly, it has been found that the limitation by a high Km, an intrinsic property of most human enzymes applicable for in vivo prodrug activation, can be overcome by novel prodrugs.

These prodrugs have the formula I,

S-Z-W (I)

25

30

35

40

45

wherein W means a pharmacologically active substance, Z stands for a self-immolative spacer or a bond and S is a moiety such that the S-Z bond is enzymatically cleaved at an at least 2-fold lower Michaelis-Menten constant compared to the natural enzyme substrate.

The prodrugs of the invention have the common characteristic to be cleaved by enzymes at significantly lower molar prodrug concentration as the natural or standard substrates used for enzymatic analysis or appropriate state of the art prodrugs (WO 92/19639). They are therefore named Km-reduced prodrugs.

The prodrugs of the invention have as another common characteristic a modified competitive enzyme activity inhibitor (S) as a crucial structural component which can be linked directly or via a spacer moiety (Z) to the pharmacologically active substance (W). Preferably the spacer is self-immolative generating the pharmacologically active substance after enzymatic cleavage of the S-Z bond. A self-immolative spacer is defined as a moiety which is bound through two bonds to two molecules and which eliminates itself from the second molecule if the bond to the first molecule is cleaved.

The preferred Km-reduced prodrugs are substrates for human glycosidases and have the general formula II:

5

10

15

$$\begin{bmatrix} R & R & R \\ R & R & CH-X-Z-W \\ R & R & R \end{bmatrix}$$
(II)

wherein

may be independent from each other H, OH, F, NH<sub>2</sub>, COOH, CH<sub>2</sub>-COOH,CHOH-COOH, PO<sub>3</sub>H<sub>2</sub>,CH<sub>2</sub>-PO<sub>3</sub>H<sub>2</sub> or CHOH-PO<sub>3</sub>H<sub>2</sub>,

X may be NH, O or S,

m may be 0 or 1,

Z stands for a self-immolative spacer or a bond and

W means a pharmacologically active substance.

Especially preferred are Km-reduced prodrugs which are substrates for  $\beta$ -glucuronidase and have the general formula III:

25

20

$$\begin{array}{c|cccc}
H & Y & X & Z - W \\
R & H & H & R
\end{array}$$
(III)

30

35

40

45

50

#### wherein

Y may be COOH, CH<sub>2</sub>-COOH, CHOH-COOH, PO<sub>3</sub>H<sub>2</sub>, CH<sub>2</sub>PO<sub>3</sub>H<sub>2</sub> or CHOH-PO<sub>3</sub>H<sub>2</sub>,

X may be NH, O or S,

R may be independent from each other F, NH2, H or OH,

m may be 0 or 1,

z stands for a bond or a self-immolative spacer preferentially a moiety with the formula

 $-Y[-C(=Y)-X-]_{p}-V(R)_{n}-X-C(=Y)-$ 

wherein

V is an aromate or a hetero aromate or an aliphate with conjugated double bonds or an amino acid residue which cycles after cleavage of the glycosyl residue, preferentially with 5-20 carbon atoms and 0-4 hetero atoms, wherein hetero atom means N, O or S, substituted with

being independently from each other H, methyl, methoxy, carboxy, methyloxycarbonyl, CN, hydroxy, nitro, fluor, chlor, brom, sulfonyl, sulfonamid or sulfon (C<sub>1-4</sub>)-alkylamid and

p 0 or 1

n an integer of 0 to 25, preferentially 1 or 2

X O, NH, methylenoxy, methylenamino or methylen  $(C_{1-4})$ -alkylamino,

O or NH

and

w means a pharmacologically active substance preferentially an anthracycline such as doxorubicin,
4'-epi-doxorubicin, 4- or 4'-desoxy-doxorubicin, or an etoposide, N-bis-(2-chlor thyl)-4-hydroxyanilin, 4-hydroxycyclophosphamide, vindesin, vinblastine, vincristine, terfenadine, terbutaline,
fenoterol, salbutamol, muscarine, oxyphenbutazone, salicylic acid, p-aminosalicylic acid, 5-

fluorouracil, 5-fluorocytidine, 5-fluorouridine, methotrexate, diclofenac, flufenamic-acid, 4-

methylaminoph nazon , theophylline, nifedipin , mitomycine C, mitoxantron , camptothecin , m-AMSA, taxol, nocodaxol, colchicine, cyclophosphamide, rachelmycin, cisplatin, melphalan, bleomycin, nitrogen-mustard, phosphoramide-mustard, quercetin, genistein, erbstatin, tyrphostin, rohitukine-d rivative ((-)-cis-5,7-dihydroxy-2-(2-chlorphenyl)-8-[4-(3-hydroxy-1-methyl)-piperidinyl]-4H-1-benzopyran-4-on; EP 89119710.5), retinoic acid, butyric acid, phorbol ester, DMSO, aclacinomycin, progesterone, buserelin, tamoxifen, mifepristone, onapristone, N-(4-aminobutyl)-5chloro-2-naphtalen-sulfonamide, pyridinyloxazol-2-one, quinolyl-, isoquinolyloxazolone-2-one, staurosporine, ethanolamine, verapamil, forskolin, 1,9-dideoxyforskolin, quinine, quinidine, reserpine, 18-O-(3,5-dimethoxy-4-hydroxybenzoyl)-reserpate, lonidamine, buthionine sulfoximine, diethyldithiocarbamate. cyclosporine A. azathioprine, chlorambucil, N-(4-trifluormethyl)-phenyl-2cyano-3-hydroxy-croton-acid-amide (WO 91/17748), 15-deoxyspergualin, FK 506, ibuprofen, indomethacin, aspirin, sulfasalazine, penicillamine, chloroquine, dexamethasone, prednisolone, lidocaine, propafenone, procaine, mefonamic acid, paracetamol, 4-aminophenazone, muskosine, orciprenaline, isoprenaline, amiloride, p-nitrophenylguanidinobenzoat or their derivatives additionly substituted with one or more hydroxy-, amino- or iminogroups, linked through a hydroxy-, aminoor imino group to Z.

not included are compounds where m=1, R=OH, Y=COOH and X=O, which is the natural substrate  $\beta$ -glucuronide. Enzyme in this application may also mean a catalytic antibody. The compounds described herein can be prepared by prior art methods.

Km-reduced prodrugs, selective for human  $\beta$ -glucuronidase, are described in the following sections. Prodrug A (example 1) may be looked upon as derived from the competitive  $\beta$ -glucuronidase inhibitor saccharolactone:

4

5

10

15

25

30

35

40

45

50

## Example 1:

Prodrug A, B, C:

10

5

NO2

ОН

20

15

25

30

Prodrug A:

R = CHOH-COOH

HO

Prodrug B:

R = CH2-COOH

OH

Prodrug C:

R = COOH

Experimental procedure for prodrug A:

Preparation of 1,2,5-tri-O-acetyl-aldehydo-D-glucurono-3,6-lactone (compound 7).

40

но

45

## Compound 5

50

3,6-Glucarolactone (compound 5) (45 g) was slowly added to a cooled (0-5  $^{\circ}$  C) mixture of dry pyridine (225 ml) and Ac<sub>2</sub>O (185 ml). The int rnal temperature was maintained at 5  $^{\circ}$  C during the addition and after all the lactone has been dissolved, the reaction mixture was allowed to be stirred for additional 2 hours. The colourless solution was then poured into 3 liters of a mixture of water and crushed ice and vigorously stirred for approximately 3 hours. The precipitate was collected and washed with water, and after drying a solid was isolated which contains 70 g of a mixture of  $\alpha$  and  $\beta$  tri-O-acetyl-glucuronolactone (compound 7). This mixture was directly used for the next step.

Preparation of 2,5-di-O-acetyl- $\alpha$ -D-glucurono-3,6-lactone- $\alpha$ -furanosyl bromide (compound 8).

Titanium bromide (16.6 g, 45 mmol) as added to a stirred solution of compound 7 (70 g, 23.3 mmol) in dichloromethane (200 ml) maintained in the dark and under nitrogen atmosphere. After stirring overnight, additional TiBr<sub>4</sub> was added (8.3, 22 mmol). After 24 additional hours, the reaction mixture was diluted with dichloromethane (150 ml) and the organic solution poured into crushed ice water. The organic layer was separated, washed with water, dried and evaporated under reduced pressure. This gave compound 8 (65 g) pure enough for the next step.

Preparation of  $(2-nitro-4-formylphenyl)-2,5-dl-O-acetyl-\beta-D-glucurono-3,6-lactone furanoside (compound 9).$ 

It was prepared from compound 8 (15 g, 50 mmol) and from 4-hydroxy-3-nitrobenzaldehyde according to the procedure already described in WO 92/ 19639. This afforded 12 g (61.6 %) of compound 9.

Preparation of  $(2-nitro-4-formylphenyl)-2,3,5-tri-O-acetyl-\beta-D-glucuronate (compound 1a).$ 

To a solution of solid sodium hydroxide (50 mg) in methanol (125 ml), compound 9 (10 g) was added.

The solution was stirred at room temperature for 4 h and evaporated under reduced pressure. This resulted in a crude mixture which was immediately dissolved in anhydrous pyridine (50 ml). After cooling to 0 ° C, acetic anhydride (40 ml) was added and the reaction mixture was subsequently stirred for additional 18 h. Extraction with dichloromethane followed by usual work-up resulted in 6.6 g of compound 1a (65 % overall yield).

Preparation of  $(2-nitro-4-hydroxymethylphenyl)-2,3,5-tri-O-acetyl-<math>\beta$ -D-glucuronate (compound 2a).

It was prepared by sodium borohydride reduction of compound 1a (6 g) according to the procedure already described in the WO 92/19639. This yielded 5.6 g (95 %) of compound 2a.

Preparation of  $4-(2,3,5-tri-O-acetyl-\beta-D-methylglucuronofuranosyl)-3-nitro-p-nitroben-zyloxycarbonate (compound 3a).$ 

# Compound 3a

It was prepared coupling of compound 2a (6 g) with 4-nitrophenyl chloroformiate (yield 75 %) according to the procedure air ady described in the WO 92/19639.

## Preparation of prodrug A:

Prodrug A was prepared from compound 3a and doxorubicin (yield 83 %) followed by treatment with sodium methoxid in methanol and then sodium hydroxide.

10

25

35

40

45

50

# Exampl 2:

Prodrug D:

5

15

20

25

30

35

45

50

55

OH OH Ō OCH<sub>3</sub> ÓН HO ÒН NO<sub>2</sub>

Experimental procedure for phenylxylopyranosylphosphonic acid (4):

1

The glucuronide 1 (with R1 = benzyl), 5 equiv. lead tetraacetate and a catalytic amount of Cu(OAc)2 in benzene are heated to reflux until the starting material has disappeared. The acetates 2 are isolated by aqueous workup (compare A.Vassella, R. Wyler, Helv. Chim. Acta 1991, 74, 451).

2

A solution of the acetat s 2 in dichlorom thane was treated with an excess of tri methylphosphate and

trimethylsilyltriflate between 0 ° and 40 ° C until the disappearance of the starting material.

3

The phosphonates 3 were isolated by aqueous workup as a mixture of diastereoisomers, which were separated by flash chromatography (compare A.Vassella, R. Wyler, Helv. Chim. Acta 1991, 74, 451).

4

The minor isomere was transformed into the phosphonic acid 4 by sequential treatment with trimethylsilyl bromide in dichloromethane and hydrogenolysis in the presence of Pd(OH)<sub>2</sub> in methanol (compare A.Vassella, R. Wyler, Helv. Chim. Acta 1991, 74, 451).

Prodrug D was synthesized analogously as described in WO 92/19639.

#### Example 3:

5

10

15

20

30

Comparison of Km- and Vmax-values of natural and improved substrate for antibody β-glucuronidase fusion protein

For Km- and Vmax-determination 3'-N-[4-(beta-D-Glucuronyloxy)-3-nitro-benzyloxycarbonyl]-doxorubicin and prodrug A were diluted in the range of 10-10000 μM in 100 mM phosphate buffer + 1 mg/ml BSA, pH 7.2. Enzymatic cleavage was done with constant amounts of fusion protein at 37 ° C. Cleavage was monitored by HPLC analysis. Km- and Vmax-values were calculated with the software program GraFit 2.0 (Erithacus Software Ltd.).

#### 40 HPLC Analysis:

The HPLC apparatus consisted of an autosampler (Abimed, model 231), an automatic sample extraction system (AASP, Varian) equipped with minicartridges containing C 18 reversed phase silica gel (Analytichem), a gradient pump (Gynkotek, model 480), a fluorescence detector (Shimazdu RF 535, Excitation: 495 nm, Emission: 560 nm). Before sample injection the minicartridges were preconditioned with 2.5 ml methanol and 1.5 ml phosphate buffer, pH 6. Analytes retained on the reversed phase silica gel were then eluted by valve switching and connection of the minicartridges to the mobile phase. Chromatography was performed on reversed phase material (Nucleosil C 18, 5 µm particle size, 120 mm length, 4,5 mm I.D.) and gradient elution. Elution was done by a gradient composed of 2 components (A: 20 mM phosphate, pH 3, B: acetonitrile). The gradient was run with the following time-concentration profile:

0 min: 75 % A, 25 % B 20 min: 25 % A, 75 % B 30 min: 25 % A, 75 % B

Befor starting the next run the column was allowed to equilibrate at starting conditions for 5 min.

55

enzym	substrat	Km mM	Vmax nmol/µg/min
antibody-8-glucuronidase fusion protein	3'-N-[4-(beta-D-glucuronyloxy)-3-nitrobe- nzyloxycarbonyl]-doxorubicin (glucuronide prodrug)	1.3	0.635
	prodrug A (Km-reduced prodrug)	< 0.05	0.71

Example 4:

Prodrug A was encapsulated according to D. Papahadjopoulos et al. (PNAS, USA 88:11460-11464, 1991) into stealth liposomes. After i.v. injection into CD1 nu/nu mice the plasma clearance of Prodrug A encapsulated into stealth liposomes was prolonged from ≈ 20 min for the free Prodrug A to ≈ 40 hrs for the encapsulated Prodrug A (data not shown). The significant t1/2β prolongation leads to improved pharmacological efficacy.

20 Claims

25

30

5

10

1. Compound according to formula I,

S-Z-W (I)

wherein

W means a pharmacologically active substance,

Z stands for a self-immolative spacer or a bond and

S is a modified competitive enzyme inhibitor such that the Z-S bond can be enzymatically cleaved at an at least 2-fold lower Michaelis Menten constant compared to the natural enzyme substrate.

- 2. Compound according to claim 1 wherein the bond between S and Z is a glycosidic bond cleavable by an enzyme or a catalytic antibody.
- 35 3. Compound according to claim 2 wherein the bond is cleavable by a human glycosidase, preferentially human β-glucuronidase.
  - 4. Compound according to claim 2 with the formula II,

45

50

55

40

$$\begin{bmatrix}
R & R & X \\
R & R & CH-X-Z-W \\
R & R & R
\end{bmatrix}$$
(II)

wherein

R may be independent from each other H, OH, F, NH<sub>2</sub>, COOH, CH<sub>2</sub>- COOH, CHOH-COOH, PO<sub>3</sub>H<sub>2</sub>, CH<sub>2</sub>-PO<sub>3</sub>H<sub>2</sub> or CHOH-PO<sub>3</sub>H<sub>2</sub>,

X may be NH, O or S,

m may be 0 or 1,

W means a pharmacologically activ substance and Z stands for a self-immolative spacer or a bond.

5. Compound according to claim 4 with the formula III,

$$\begin{bmatrix}
H & Y & O & Z & W \\
H & R & H & H
\end{bmatrix}$$
(III)

wherein

5

10

15

20

25

35

40

45

50

55

may be COOH, CH2-COOH, CHOH-COOH, PO3H2, CH2PO3H2 or CHOH-PO3H2. Υ

X may be NH, O or S.

R may be independent from each other F, NH2, H or OH,

m may be 0 or 1,

stands for a bond or a self-immolative spacer preferentially a moiety with the formula Ζ

 $-Y[-C(=Y)-X-]_{0}-V(R)_{0}-X-C(=Y)-$ 

wherein

is an aromate or a hetero aromate or an aliphate with conjugated double bonds or an amino acid residue which cycles after cleavage of the glycosyl residue, preferentially with 5-20 carbon atoms and 0-4 hetero atoms, wherein hetero atom means N, O or S, substituted with

being independently from each other H, methyl, methoxy, carboxy, methyloxycarbonyl, CN, R hydroxy, nitro, fluor, chlor, brom, sulfonyl, sulfonamid or sulfon (C1-4)-alkylamid and

р

an integer of 0 to 25, preferentially 1 or 2 n

O, NH, methylenoxy, methylenamino or methylen (C1-4)-alkylamino, Х

O or NH

30 and

> means a pharmacologically active substance, preferentially an anthracycline such as doxorubicin, 4'-epi-doxorubicin, 4- or 4'-desoxy-doxorubicin, or an etoposide, N-bis-(2-chlorethyl)-4-hydroxyaniline, 4-hydroxycyclophosphamide, vindesine, vinblastine, vincristine, terfenadine, terbutaline, fenoterol, salbutamol, muscarine, oxyphenbutazone, salicylic acid, p-aminosalicylic acid, 5-fluorouracil, 5-fluorocytidine, 5- fluorouridine, methotrexate, diclofenac, flufenamicacid, 4-methylaminophenazone, theophylline, nifedipine, mitomycine C, mitoxantrone, camptothecine, m-AMSA, taxol, nocodaxol, colchicine, cyclophosphamide, rachelmycin, cisplatin, melphalan, bleomycin, nitrogen-mustard, phosphoramide-mustard, quercetln, genistein, erbstatin, tyrphostin, rohitukine-derivative ((-)-cis-5,7-dihydroxy-2-(2-chlorphenyl)-8-[4-(3-hydroxy-1-methyl)-piperidinyl]-4H-1-benzopyran-4-on; EP 89119710.5), retinoic acid, butyric acid, phorbol ester, DMSO, aclacinomycin, progesterone, buserelin, tamoxifen, mifepristone, onapristone, N-(4-aminobutyl)-5-chloro-2-naphtalen-sulfonamide, pyridinyloxazol-2-one, quinolyl-, isoquinolyloxazolone-2-one, staurosporine, ethanolamine, verapamil, forskolin, 1,9-dideoxyforquinine, quinidine, reserpine, 18-O-(3,5-dimethoxy-4-hydroxybenzoyl)-reserpate, lonidamine, buthionine sulfoximine, diethyldithiocarbamate, cyclosporine A, azathioprine, chlorambucil, N-(4-trifluormethyl)-phenyl-2-cyano-3-hydroxy-croton-acid-amide 91/17748), 15-deoxyspergualin, FK 506, ibuprofen, indomethacin, aspirin, sulfasalazine, penicillamine, chloroquine, dexamethasone, prednisolone, lidocaine, propafenone, procaine, mefonamic acid, paracetamol, 4-aminophenazone, muskosine, orciprenaline, isoprenaline, amiloride, p-nitrophenylguanidinobenzoat or their derivatives additionly substituted with one or more hydroxy-, amino- or iminogroups, linked through a hydroxy-, amino- or imino group to Z. Not included are compounds where m = 1, R = OH, Y = COOH and X = O.

A pharmaceutical containing a compound according to claim 1.

A pharmaceutical according to claim 5 encapsulated in liposomes.

8. A pharmaceutical according to claim 5 in combination with pretargeted enzymes, catalytic antibodies, immunotoxins or immunoconjugates. 



# PARTIAL EUROPEAN SEARCH REPORT

Application Number

which under Rule 45 of the European Patent Convention EP 93 11 4475 shall be considered, for the purposes of subsequent proceedings, as the European search report

DOCUMENTS CONSIDERED TO BE RELEVANT		İ		
Category	Citation of document wit of relevant	n indication, where appropriate, passages	Reirvant to claim	CLASSIFICATION OF THE APPLICATION (Int.CL6)
D,X, Y	EP-A-0 511 917 (L	AB. HOECHST S.A.)	1-7	A61K47/48
	* page 5, line 1	1ine 50 *		
	* page 6, line 25; & WO-A-92 19639	claims *		
<b>(,</b> Y	EP-A-O 501 215 (BE * example T *	HRINGWERKE AG.)	1-7	
<b>'</b>	WO-A-81 01145 (UNI FOUNDATION)	VERSITY OF ILLINOIS	1-7	
	* page 19; figure	I_*	1	
	* page 21; table 2 * page 34, line 23	; claims *		
,	WO-A-90 03188 (NEO	RX CO.)	1-7	
	* page 19, line 27	- page 24, line 33 *		
	EP-A-O 540 859 (BR * claim 1 *	ISTOL-MYERS SQUIBB CO.)	1-7	
.	EP-A-0 441 218 (BE	HRINGWERKE)	1-7	TECHNICAL FIELDS SEARCHED (Lt.Cl.6)
	* page 5; claims;	figures I,ÍI *		A61K
NCO	MPLETE SEARCH		<u> </u>	
ot a mean laims sea laims sea laims act	a Division considers that the presents of the European Patent Conventaging search into the state of the arched completely: school incompletely: searched; the limitation of the search:	European patent application does not comply tion to such an extent that it is not possible to rt on the basis of some of the claims	with curry	
	Shoot C			·
500	meet t			
see s				
	Place of search	Date of completion of the search		Domine
7	HE HAGUE	14 March 1994	Bert	<del>Pomini</del> e, M
CA  C : partice   . —	14 March 1994  T: theory or principle E: earlier patent éous	underlying the it ment, but publishe the application other reasons	ee, M ovention hed as, or	

EPO PORM 1500 CC.62 (POLCOT)



EP 93 11 4475

-C-

INCOMPLETE SEARCH

Claims searched incompletely: 1-7

In view of the large number of compounds, which are designed by the general formulas of claim 1 and also in view of the definition of products by means of their biological, chemical and/or pharmacological properties, the search has to be restricted for economic reasons. The search was limited to the compounds for which pharmacological data was given and/or the compounds mentioned in the claims or examples (see Guidelines, Part B, Chapter III, paragraph 3.6)

1.0